

Tue Jul 17 13:47:10 2001 [BLASTN 2.1.3 [Apr-1-2001], NCBI] $^{\prime}$ home/ruby/va/Molbio/carpenda/templ/ss.DNA32292 (1364 bp)

1 P_AAC974 2 P_AAF723 3 P_AAF603 4 P_AAZ937 5 P_AAA300 6 P_AAA540 7 P_AAX284 8 P_AAX522 9 P_AAX376 10 AX076909 11 P_AAF294 12 P_AAA085 13 P_AAC769 >1 P_AAC9740 (1364 bp) [1 Score = 136	7 Human TANGO 331 cDNA. + 1350 1350 100 0. 8 DNA encoding human apoptosis related pro + 1290 1358 99 0. 9 Human ORFX ORF2475 polynucleotide sequen + 1208 1215 100 0. 9 Human angiogenesis-associated protein PRO211 cDNA, SEQ ID NO:56.	0 0 0 0 0 0 0 0 0
DNA32292	1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC	
P AAC97409	**************************************	
_		
DNA32292	61 CGCCATGCGCCTGCCGCGCCGGGCCGCGCTTCTGCCGCTTCTGCTG	
P_AAC97409	61 CGCCATGCGCCTGCCGCCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG	
DNA32292	21 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA	
P_AAC97409	.21 GCCCGCGCCGGAGGCCCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA	
DNA32292	81 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC	
P_AAC97409	81 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC	
DNA32292	41 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT	
P_AAC97409	**************************************	
DNA32292	01 GGAGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA	
P_AAC97409	01 GGAGGGGCTGTGCGAGAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA	
DNA32292	61 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT	
P_AAC97409	61 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT	
DNA32292	21 TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC	
P_AAC97409	21 TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC	
DNA32292	81 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGAATGGCCACTGCAGCGGAGATGGGAG	

P_AAC97409	481	**************************************
DNA32292	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA ************************************
P_AAC97409	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAC97409	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT ****************************
P_AAC97409	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC *************************
P_AAC97409	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG *******************************
P_AAC97409	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAC97409	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
P_AAC97409	901	
DNA32292	961	AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAC97409	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC *******************************
P_AAC97409	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
P_AAC97409	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAC97409	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAC97409	1201	GCAGTGGACAGCGGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT *********************************
P_AAC97409	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA

>2 P AAF72371 Human PRO211 cDNA. (1825 bp) [1 seg] Score = 1364 (2704 bits), Expect = 0.0Identities = 1364/1364 (100%), at 1,75-1364,1438, Strand +/+ DNA32292 1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC ***************** P AAF72371 75 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC DNA32292 *********************** P AAF72371 121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA DNA32292 ****************** P AAF72371 195 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA DNA32292 181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC ******************* 255 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC P AAF72371 DNA32292 241 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT ************************ P AAF72371 315 TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT 301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA DNA32292 ******************** P AAF72371 375 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA DNA32292 361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT ***************** P AAF72371 435 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT DNA32292 421 TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC ****************** P AAF72371 495 TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC 481 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG DNA32292 ***************** P AAF72371 555 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG DNA32292 541 CAGACAGGGCGACGGTTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA ******************** P AAF72371 615 CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA DNA32292 601 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG ******************* P AAF72371 675 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG DNA32292 661 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT P AAF72371 735 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT DNA32292 721 GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC ***** P AAF72371 795 GGGCTGGGTGCTGGACGAGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC

DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG		
P_AAF72371	855	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG		
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC *********************************		
P_AAF72371	915	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAAGGCCCAGGAAACTGTAAAGAGTGTATCTC		
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA		
P_AAF72371	975	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA		
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG		
P_AAF72371	1035	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG		
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC *******************************		
P_AAF72371	1095	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC		
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC *********************************		
P_AAF72371	1155	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC		
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT		
P_AAF72371	1215	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT		
DNA32292	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC		
P_AAF72371	1275	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC		
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT *********************************		
P_AAF72371	1335	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT		
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA		
P_AAF72371	1395	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA		
>3 P_AAF60360 PRO211 coding sequence. (1364 bp) [1 seg] Score = 1364 (2704 bits), Expect = 0.0 Identities = 1364/1364 (100%), at 1,1-1364,1364, Strand +/+				
DNA32292				
		GGCCGGAGCACGCCGCAGGACCTGGAGCTCCGGCTGCTTTCCCGCAGCGCTACC **********************************		
P_AAF60360		GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC		
DNA32292		CGCCATGCGCCTGCCGGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG		
P_AAF60360	61	CGCCATGCGCCTGCCGGGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG		
DNA32292	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGCTGGTGGA ********************************		
P_AAF60360	121	GCCCGCGCGGAGGCCGCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA		

DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
P_AAF60360	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT **********************************
P_AAF60360	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA ************************
P_AAF60360	301	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT *******************************
P_AAF60360	361	GCACCTGGAGGCCTGGTGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC ******************************
P_AAF60360	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG ****************************
P_AAF60360	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA ************************************
P_AAF60360	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG **********************************
P_AAF60360	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT ****************************
P_AAF60360	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC *************************
P_AAF60360	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG *******************************
P_AAF60360	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC *********************************
P_AAF60360	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA ******************************
P_AAF60360	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAF60360	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC

P_AAF60360	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
P_AAF60360	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAF60360	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAF60360	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
P_AAF60360	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
P_AAF60360	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
Score = 136	54 (2	0211 DNA32292-1131. (1364 bp) [1 seg] 704 bits), Expect = 0.0 64/1364 (100%), at 1,1-1364,1364, Strand +/+
DNA32292	1	GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
P_AAZ93700	1	GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
DNA32292	61	CGCCATGCGCCTGCCGCGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG
P_AAZ93700	61	CGCCATGCGCCTGCCGCGCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
P_AAZ93700	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
P_AAZ93700	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
P_AAZ93700	241	TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
P_AAZ93700	301	GGAGGGGCTGTGCGAGGCGACCTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
P_AAZ93700	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC

P_AAZ93700	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
P_AAZ93700	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
P_AAZ93700	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAZ93700	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
P_AAZ93700	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAZ93700	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
P_AAZ93700	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAZ93700	841	
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
P_AAZ93700	901	TGGCTACGCGAGGGACACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGT
P_AAZ93700	961	AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
P_AAZ93700	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
P_AAZ93700	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAZ93700	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAZ93700	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
P_AAZ93700	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT

```
DNA32292
         P AAZ93700
         >5 P_AAA30040 Human PRO211 nucleotide sequence. (1364 bp) [1 seg]
 Score = 1364 (2704 bits), Expect = 0.0
 Identities = 1364/1364 (100%), at 1,1-1364,1364, Strand +/+
           1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
 DNA32292
             *****************
           1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
P AAA30040
 DNA32292
          *****************
P AAA30040
          DNA32292
         121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
             *************
P AAA30040
         121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
 DNA32292
         181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
             *****************
P AAA30040
         181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
 DNA32292
         241 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
P AAA30040
         241 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
 DNA32292
         301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
             ***********
P AAA30040
         301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
 DNA32292
         361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
            *******************
P AAA30040
         361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
         421 TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC
 DNA32292
P AAA30040
         421 TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
 DNA32292
         481 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
            ******************
P AAA30040
         481 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
 DNA32292
         541 CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
            ***********************
P AAA30040
         541 CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
 DNA32292
         601 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
            P AAA30040
         601 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
 DNA32292
         661 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
            **********
P AAA30040
         661 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
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DNA32292	721	GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAA30040	721	GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
P_AAA30040	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAA30040	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGACCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
P_AAA30040	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAA30040	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGT
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
P_AAA30040	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
P_AAA30040	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAA30040	-1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAA30040	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
P_AAA30040	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
P_AAA30040	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
Score = 13	64 (27	D211 cDNA. (1364 bp) [1 seg] 704 bits), Expect = 0.0 64/1364 (100%), at 1,1-1364,1364, Strand +/+
DNA32292	1	GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
P_AAA54089	1	GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
DNA32292	61	CGCCATGCGCCTGCCGCCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
P_AAA54089	61	CGCCATGCGCCTGCCGGGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA

P_AAA54089	121	**************************************
DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
P_AAA54089	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
P_AAA54089	241	TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
P_AAA54089	301	GGAGGGGCTGTGCGAGGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
P_AAA54089	361	GCACCTGGAGGCCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
P_AAA54089	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
P_AAA54089	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
P_AAA54089	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAA54089	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
P_AAA54089	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAA54089	721	GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
P_AAA54089	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAA54089	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
P_AAA54089	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG

P_AAA54089	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
P_AAA54089	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC *********************************
P_AAA54089	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAA54089	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAA54089	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
P_AAA54089	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
P_AAA54089	1321	**************************************
(1364 bp) [Score = 13	1 seg] 64 (2)	F-like homologue PRO211 coding sequence. DNA, PAT 22-JUN-1999 704 bits), Expect = 0.0 64/1364 (100%), at 1,1-1364,1364, Strand +/+
DNA32292		GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
P AAX28433		**************************************
DNA32292		CGCCATGCGCCTGCCGCGCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
P AAX28433		*****************
		CGCCATGCGCCTGCCGCGCCGGGCCGCGCTGGGGGCTCCTGCCGCTTCTGCTG
DNA32292		GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA ********************************
P_AAX28433	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292		
P_AAX28433	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC **********************
		CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGAACACGGC ***********************
DNA32292	181	**************
DNA32292 P_AAX28433	181 241	**************************************
	181 241 241	**************************************
P_AAX28433	181 241 241 301	**************************************

P_AAX28433	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC ******************************
P_AAX28433	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
P_AAX28433	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
P_AAX28433	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAX28433	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT ****************************
P_AAX28433	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAX28433	721	GGGCTGGGTGCTGGACGAGGGCCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG *******************************
P_AAX28433	781	$\tt TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG$
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAX28433	841	TGACTCCAGCTGTGGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA ******************************
P_AAX28433	901	$\tt TGGCTACGCGAGGGGACACGGACAGTGTGCAGAGGAGGGGGGGG$
DNA32292	961	AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAX28433	961	${\tt AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG$
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
P_AAX28433	1021	${\tt TCCTGACGGCTTCGAAGAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC}$
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
P_AAX28433	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAX28433	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAX28433	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC

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1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
 DNA32292
            *****************
P AAX28433
         1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
 DNA32292
        ***************
P AAX28433
        >8 P AAX52213 Protein PRO211 cDNA clone DNA32292-1131. DNA, PAT 25-JUN-1999
(1364 bp) [1 seq]
 Score = 1364 (2704 bits), Expect = 0.0
 Identities = 1364/1364 (100%), at 1,1-1364,1364, Strand +/+
 DNA32292
           1 GGCCGGAGCACGCCGCAGGACCTGGAGCTCCGGCTGCTCTTCCCGCAGCGCTACC
            ******************
           P AAX52213
 DNA32292
          P AAX52213
          DNA32292
         121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
            *****************
P AAX52213
         121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
 DNA32292
         181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
            ******************
P AAX52213
         181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
 DNA32292
         241 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
            **********************
P AAX52213
         241 TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
 DNA32292
         301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
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P AAX52213
         301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
 DNA32292
         361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
            ******************
P AAX52213
         361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
         421 TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
 DNA32292
            P AAX52213
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 DNA32292
         481 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
            *****************
P AAX52213
         481 ATGCCAGGGGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
 DNA32292
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            P AAX52213
         541 CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
         601 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG
 DNA32292
            P AAX52213
         601 CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG
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661 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
   DNA32292
              *******************
           661 TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
 P AAX52213
           721 GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
  DNA32292
              **********************
           721 GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
 P AAX52213
           781 TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
  DNA32292
              **********************
 P AAX52213
           781 TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
           841 TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
  DNA32292
              ***********************
 P AAX52213
          841 TGACTCCAGCTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
          901 TGGCTACGCGAGGGGGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
  DNA32292
              *******************
 P AAX:52213
          901 TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
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              **********************
          961 AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG
P AAX52213
         1021 TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
  DNA32292
             ******************
P AAX52213
         1021 TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
  DNA32292
         1081 AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
             ************************
P AAX52213
         1081 AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
         1141 CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
  DNA32292
             ***********************
P AAX52213
         1141 CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
 DNA32292
         1201 GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTAACGGTTGATTCTCATTTGTCCC
             ***********************
P AAX52213
         1201 GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
         1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
 DNA32292
             **********************
         1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
P AAX52213
 DNA32292
         ****************
P AAX52213
         >9 P_AAX37671 Human EGF-like homologue (PRO217) encoded by DNA32292 cDNA. (1364
bp) [1 sea]
Score = 1364 (2704 bits), Expect = 0.0
Identities = 1364/1364 (100%), at 1,1-1364,1364, Strand +/+
           DNA32292
             *******************
P AAX37671
           1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
```

DNA32292	61	CGCCATGCGCCTGCCGCGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG
P_AAX37671	61	CGCCATGCGCCTGCCGCGCCGGGCCGCGCTGGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121	GCCCGCGCGGGGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA ********************************
P_AAX37671	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC **********************
P_AAX37671	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT **********************************
P_AAX37671	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA ************************
P_AAX37671	301	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	361	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT *******************************
P_AAX37671	361	GCACCTGGAGGCCTGGTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC ******************************
P_AAX37671	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGAATGGCCACTGCAGCGGAGATGGGAG ****************************
P_AAX37671	481	ATGCCAGGGGGATCCCAGAGGCCCTGCAGCGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
P_AAX37671	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAX37671	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGCCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT ****************************
P_AAX37671	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAX37671	721	GGGCTGGGTGCTGGACGAGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG *******************************
P_AAX37671	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAX37671	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCCAGGAAACTGTAAAGAGTGTATCTC

DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAA
P_AAX37671	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAX37671	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG
DNA32292	1021	TCCTGACGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC *******************************
P_AAX37671	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC *********************************
P_AAX37671	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAX37671	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAX37671	1201	GCAGTGGACAGCGGCGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT *********************************
P_AAX37671	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
P_AAX37671	1321	TGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
>10 AX07690	9 Sequ	uence 21 from Patent W00105836. (1364 bp) [1 seq]
Score = 13	64 (27	704 bits), Expect = 0.0 64/1364 (100%), at 1,1-1364,1364, Strand +/+
DNA32292		GGCCGGAGCACGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC **********************************
AX076909	1	GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
DNA32292	61.	CGCCATGCGCCTGCCGCCGGGCCGCGGGCTGGGGCTCCTGCCGCTTCTGCTG
AX076909	61	CGCCATGCGCCTGCCGGCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121	GCCCGCGCGGGGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA ********************************
AX076909	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
AX076909	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT **********************************
AX076909		TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA

AX076909	30:	1 GGAGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	363	GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
AX076909	361	1 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
AX076909	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
AX076909	481	ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
AX076909	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
AX076909	601	**************************************
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
AX076909	661	**************************************
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCCCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
AX076909	721	**************************************
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
AX076909	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
AX076909	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
AX076909	901	TGGCTACGCGAGGGACACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
AX076909	961	AACCTGTGTGAGGAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
AX076909	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
AX076909		AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT

AX076909	1141 CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201 GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
AX076909	1201 GCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT *********************************
AX076909	1261 TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321 TGTAATAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
AX076909	1321 TGTAATAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA
Score = 13	457 Human TANGO 331 cDNA. (1432 bp) [1 seg] 50 (2676 bits), Expect = 0.0 = 1350/1350 (100%), at 1,50-1350,1399, Strand +/+
DNA32292	1 GGCCGGAGCACGGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC **********************************
P_AAF29457	50 GGCCGGAGCACGCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
DNA32292	61 CGCCATGCGCCTGCCGCGCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
P_AAF29457	110 CGCCATGCGCCTGCCGCGCCGGGCCGCTGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121 GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA ********************************
P_AAF29457	170 GCCCGCGCCGGAGGCCCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292	181 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
P_AAF29457	230 CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241 TTGGGAGGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT **********************************
P_AAF29457	290 TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
P_AAF29457	350 GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292	361 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT *******************************
P_AAF29457	410 GCACCTGGAGGCCTGGTGGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421 TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC
P_AAF29457	470 TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG ****************************
P_AAF29457	530 ATGCCAGGGCGGATCCCAGAGGCCCTGCAGCGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541 CAGACAGGGCGACGGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
P_AAF29457	**************************************

DNA32292	601	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
P_AAF29457	650	CTGCATGGACGGCTACTTCAGCTCGCTCCGGAACGAGACCCACAGCATCTGCACAGCCTG
DNA32292	661	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
P_AAF29457	710	TGACGAGTCCTGCAAGACGTGCTCGGGCCTGACCAACAGAGACTGCGGCGAGTGTGAAGT
DNA32292	721	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
P_AAF29457	770	GGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAGCCGCC
DNA32292	781	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
P_AAF29457	830	TCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAAGAGTG
DNA32292	841	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
P_AAF29457	890	TGACTCCAGCTGTGTGGGCTGCACAGGGGAAAGGCCCAGGAAACTGTAAAGAGTGTATCTC
DNA32292	901	TGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
P_AAF29457	950	TGGCTACGCGAGGGACACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCAGAAAA
DNA32292	961	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTGTG
P_AAF29457	1010	AACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGTGTG
DNA32292	1021	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
P_AAF29457	1070	TCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAAGCCAC
DNA32292	1081	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC *********************************
P_AAF29457	1130	AGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGACTTAC
DNA32292	1141	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
P_AAF29457	1190	CCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGTCTCCT
DNA32292	1201	GCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTTGTCCC
P_AAF29457	1250	GCAGTGGACAGCGGCGGGAGAGGCTGCCTCTCTAACGGTTGATTCTCATTTGTCCC
DNA32292	1261	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT *********************************
P_AAF29457	1310	TTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAGTTCTT
DNA32292	1321	TGTAATAAATTGACCATTGTAGGTAATCA
P_AAF29457	1370	TGTAATAAAATTGACCATTGTAGGTAATCA
>12 P_AAA089 Score = 129	503 DN 90 (25	A encoding human apoptosis related protein. (1380 bp) [1 seg] 57 bits), Expect = 0.0

>12 P_AAA08503 DNA encoding human apoptosis related protein. (1380 bp) [1 segs Score = 1290 (2557 bits), Expect = 0.0 Identities = 1358/1369 (99%), Gaps = 10/1369 (0%), at 1,10-1364,1373, Strand +/+

DNA32292	1	GGCCGGAGCACGCCCGCAGGACCTGGAGCTCCGGCTGCGTCTTCCCGCAGCGCTACC
P_AAA08503	10	
DNA32292	61	CGCCATGCGCCTGCCGCCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
P_AAA08503	70	CGCCATGCGCCTGCCGCGCGGGCCGCGCTGGGGCTCCTGCCGCTTCTGCTG
DNA32292	121	GCCCGCGCGGAGGCCGCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGCTGGTGGA ********************************
P_AAA08503	130	GCCCGCGCGGAGGCCCCAAGAAGCCGACGCCCTGCCACCGGTGCCGGGGGCTGGTGGA
DNA32292	181	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC **********************
P_AAA08503	190	CAAGTTTAACCAGGGGATGGTGGACACCGCAAAGAAGAACTTTGGCGGCGGGAACACGGC
DNA32292	241	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT **********************************
P_AAA08503	250	TTGGGAGAAAAGACGCTGTCCAAGTACGAGTCCAGCGAGATTCGCCTGCTGGAGATCCT
DNA32292	301	GGAGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA ************************
P_AAA08503	310	GGAGGGGCTGTGCGAGAGCAGCGACTTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
DNA32292		GCACCTGGAGGCCTGGTGCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT *******************************
P_AAA08503		GCACCTGGAGGCCTGCAGCTGAAGAGCGAATATCCTGACTTATTCGAGTGGTT
DNA32292	421	TTGTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCCGACTGTCTCGC ******************************
P_AAA08503	430	TTGTGTGAAGACACTGAAAGTGTGCTGCTCCCAGGAACCTACGGTCCCGACTGTCTCGC
DNA32292	481	ATGCCAGGGGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG ****************************
P_AAA08503		ATGCCAGGGGGATCCCAGAGGCCCTGCAGCGGGAATGGCCACTGCAGCGGAGATGGGAG
DNA32292	541	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA ************************************
P_AAA08503	550	CAGACAGGGCGACGGTCCTGCCGGTGCCACATGGGGTACCAGGGCCCGCTGTGCACTGA
DNA32292	601	CTGCATGGACGGCTACTTCAGCT-CGCTCCGG-AACGAGACCCACAGCATCTGCACAGCC *******************************
P_AAA08503		CTGCATGGACGGCTACTTCAGCTCCGCTCCGGRAACGAGACCCACAGCATCTGCACAGCC
DNA32292		TGTGACGAGTCCTGCAAGACGTGCTC-GGGCCT-GACCAACAGAGAC-TGCGGCGAGTGT *********************************
P_AAA08503		TGTGACGAGTCCTGCAAGACGTGCTCGGGGCCTGGACCAACAGAGACTTGCGGCGAGTGT
DNA32292		GAAGTGGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGAG ******************************
P_AAA08503		GAAGTGGGCTGGGTGCTGGACGAGGGCGCCTGTGTGGATGTGGACGAGTGTGCGGCCGA-
DNA32292		CCGCCTCCCTGCAGCGCTGCGCAGTTCTGTAAGAACGCCAACGGCTCCTACACGTGCGAA *** ******** *** *******************
P_AAA08503	789	CCGTCCCTGCAGCG-TGC-CAGTTCTGTAAGAACCCCAACGGCTCCTACACGTGCGAA

DNA32292	836 GAGTGTGACTCCAGCTGTGTGGGCTGCACAGGGGAAGGCCCAGGAAACTGTAAAGAGTGT ***************************	
P_AAA08503	845 GAGTGTGACTCCAGCTGTGTGGGCTGCACAGGGGAAAGGCCCCAGGAAACTGTAAAGAGTGT	
DNA32292	896 ATCTCTGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCA ***********************************	
P_AAA08503	905 ATCTCTGGCTACGCGAGGGAGCACGGACAGTGTGCAGATGTGGACGAGTGCTCACTAGCA	
DNA32292	956 GAAAAAACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGT ********************************	(
P_AAA08503	965 GAAAAAACCTGTGTGAGGAAAAACGAAAACTGCTACAATACTCCAGGGAGCTACGTCTGT	
DNA32292	O16 GTGTGTCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAA	
P_AAA08503	.025 GTGTGTCCTGACGGCTTCGAAGAAACGGAAGATGCCTGTGTGCCGCCGGCAGAGGCTGAA	(
DNA32292	.076 GCCACAGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGA	
P_AAA08503	085 GCCACAGAAGGAGAAAGCCCGACACAGCTGCCCTCCCGCGAAGACCTGTAATGTGCCGGA	
DNA32292	136 CTTACCCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGT	
P_AAA08503	145 CTTACCCTTTAAATTATTCAGAAGGATGTCCCGTGGAAAATGTGGCCCTGAGGATGCCGT	
DNA32292	196 CTCCTGCAGTGGACAGCGGCGGGAGAGGCTGCCTGCTCTCTAACGGTTGATTCTCATTT *****************************	
P_AAA08503	205 CTCCTGCAGTGGACAGCGGCGGGGAGAGGCTGCCTGCTCTAACGGTTGATTCTCATTT	
DNA32292	256 GTCCCTTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAG	
P_AAA08503	265 GTCCCTTAAACAGCTGCATTTCTTGGTTGTTCTTAAACAGACTTGTATATTTTGATACAG	
DNA32292	316 TTCTTTGTAATAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA	
P_AAA08503	325 TTCTTTGTAATAAAATTGACCATTGTAGGTAATCAGGAGGAAAAAAAA	
pp) [1 seg]	O Human ORFX ORF2475 polynucleotide sequence SEQ ID NO:4949. (1259	
Score = 120 Identities +/+	(2395 bits), Expect = 0.0 $1215/1216 (99%)$, Gaps = $1/1216 (0%)$, at $150,32-1364,1247$, Strand	
DNA32292	150 CGCCCTGCCACCGGTGCCGGGGGCTGGTGGACAGTTTAACCAGGGGATGGTGGACACCG	
P_AAC76920	32 CGCCCTGCCACCGGTGCCGGGGGCTGGTGGACAAGTTTAACCAGGGGATGGTGGACACCG	
DNA32292	210 CAAAGAAGAACTTTGGCGGCGGGAACACGGCTTGGGAGGAAAAGACGCTGTCCAAGTACG ************************************	
P_AAC76920	92 CAAAGAAGAACTTTGGCGGCGGGAACACGGCTTGGGAGGAAAAGACGCTGTCCAAGTACG	
DNA32292	270 AGTCCAGCGAGATTCGCCTGGAGAGTCCTGGAGGGGCTGTGCGAGAGCAGCGACTTCG **********************************	
P_AAC76920	152 AGTCCAGCGAGATTCGCCTGCAGAGATCCTGGAGGGGCTGTGCGAGAGCAGCGACTTCG	
DNA32292	AATGCAATCAGATGCTAGAGGCGCAGGAGGAGCACCTGGAGGCCTGGTGGCTGCAGCTGA	
P_AAC76920	212 AATGCAATCAGATGCTAGAGGCGCAGGAGGAGCACCTGGAGGCCTGGTGGCTGCAGCTGA	

DNA32292	390	AGAGCGAATATCCTGACTTATTCGAGTGGTTTTGTGTGAAGACACTGAAAGTGTGCTGCT
P_AAC76920	272	AGAGCGAATATCCTGACTTATTCGAGTGGTTTTTGTGTGAAGACACTGAAAGTGTGCTGCT
DNA32292	450	CTCCAGGAACCTACGGTCCCGACTGTCTCGCATGCCAGGGCGGATCCCAGAGGCCCTGCA
P_AAC76920	332	CTCCAGGAACCTACGGTCCCGACTGTCTCGCATGCCAGGGCGGATCCCAGAGGCCCTGCA
DNA32292	510	GCGGGAATGGCCACTGCAGCGGAGATGGGAGCAGACAGGGCGACGGGTCCTGCCGGTGCC
P_AAC76920	392	GCGGGAATGGCCACTGCAGCGGAGATGGGAGCAGACAGGGCGACGGGTCCTGCCGGTGCC
DNA32292	570	ACATGGGGTACCAGGGCCCGCTGTGCACTGACTGCATGGACGGCTACTTCAGCTCGCTC
P_AAC76920	452	ACATGGGGTACCAGGGCCCGCTGTGCACTGACTGCATGGACGGCTACTTCAGCTCGCTC
DNA32292	630	GGAACGAGACCCACAGCATCTGCACAGCCTGTGACGAGTCCTGCAAGACGTGCTCGGGCC
P_AAC76920	512	GGAACGAGACCCACAGCATCTGCACAGCCTGTGACGAGTCCTGCAAGACGTGCTCGGGCC
DNA32292	690	TGACCAACAGAGACTGCGGCGAGTGTGAAGTGGGCTGGGTGCTGGACGAGGGCGCCTGTG
P_AAC76920	572	TGACCAACAGAGACTGCGGCGAGTGTGAAGTGGGCTGGGTGCTGGACGAGGGCCCCTGTG
DNA32292	750	TGGATGTGGACGAGTGTGCGGCCGAGCCGCCTCCCTGCAGCGCTGCGCAGTTCTGTAAGA ********************************
P_AAC76920	632	TGGATGTGGACGAGTGTGCGGCCGAGCCGCCTCCCTGCAGCGCTGCGCAGTTCTGTAAGA
DNA32292	810	ACGCCAACGGCTCCTACACGTGCGAAGAGTGTGACTCCAGCTGTGTGGGCTGCACAGGGG
P_AAC76920	692	ACGCCAACGGCTCCTACACGTGCGAAGAGTGTGACTCCAGCTGTGTGGGCTGCACAGGGG
DNA32292	870	AAGGCCCAGGAAACTGTAAAGAGTGTATCTCTGGCTACGCGAGGGAGCACGGACAGTGTG *********************************
P_AAC76920	752	AAGGCCCAGGAAACTGTAAAGAGTGTATCTCTGGCTACGCGAGGGAGCACGGACAGTGTG
DNA32292	930	CAGATGTGGACGAGTGCTCACTAGCAGAAAAACCTGTGTGAGGAAAAACGAAAACTGCT
P_AAC76920	812	CAGATGTGGACGAGTGCTCACTAGCAGAAAAACCTGTGTGAGGAAAAACGAAAACTGCT
DNA32292	990	ACAATACTCCAGGGAGCTACGTCTGTGTGTCCTGACGGCTTCGAAGAA-ACGGAAGAT ********************************
P_AAC76920	872	ACAATACTCCAGGGAGCTACGTCTGTGTGTCCTGACGGCTTCGAAGAANACGGAAGAT
DNA32292	1049	GCCTGTGTGCCGCCGGCAGAGGCTGAAGCCACAGAAGGAGAAAGCCCGACACAGCTGCCC
P_AAC76920	932	GCCTGTGTGCCGCCGGCAGAGGCTGAAGCCACAGAAGGAGAAAGCCCGACACAGCTGCCC
DNA32292	1109	TCCCGCGAAGACCTGTAATGTGCCGGACTTACCCTTTAAATTATTCAGAAGGATGTCCCG
P_AAC76920	992	TCCCGCGAAGACCTGTAATGTGCCGGACTTACCCTTTAAATTATTCAGAAGGATGTCCCG
DNA32292	1169	TGGAAAATGTGGCCCTGAGGATGCCGTCTCCTGCAGTGGACAGCGGCGGGGAGAGGCTGC
P_AAC76920		TGGAAAATGTGGCCCTGAGGATGCCGTCTCCTGCAGTGGACAGCGGCGGGGAGAGGCTGC

DNA32292	1229 CTGCTCTAACGGTTGATTCTCATTTGTCCCTTAAACAGCTGCATTTCTTGGTTGTTCT
P_AAC76920	1112 CTGCTCTCTAACGGTTGATTCTCATTTGTCCCTTAAACAGCTGCATTTCTTGGTTGTTCT
DNA32292	1289 TAAACAGACTTGTATATTTTGATACAGTTCTTTGTAATAAATTGACCATTGTAGGTAAT
P_AAC76920	1172 TAAACAGACTTGTATATTTTGATACAGTTCTTTGTAATAAAATTGACCATTGTAGGTAAT
DNA32292	1349 CAGGAGGAAAAAAA *********
P_AAC76920	1232 CAGGAGGAAAAAAA



Tue Jul 17 13:23:35 2001 [BLASTP 2.1.3 [Apr-1-2001], NCBI] /home/ruby/va/Molbio/carpenda/temp1/p1.DNA32292.nc (353 aa)

Sequences pro 1 P_AAB5307 2 P_AAB6123 3 P_AAB8021 4 P_AAB6859 5 P_AAY8322 6 P_AAB0016 7 P_AAY0528 8 P_AAY1334 9 P_AAY0806 10 P_AAY8857 11 P_AAB6123 12 AAH02894. 13 NP_077300 14 P_AAB4271 15 P_AAY9187	75 31 12 96 24 59 33 14 54 71 33 114	Human angiogenesis-associated protein PRO 2005 353 100 0.0 Human TANGO 331 protein - Homo sapiens. 2005 353 100 0.0 PRO211 protein - Homo sapiens. 2005 353 100 0.0 PRO211 Polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 Polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 polypeptide - Homo sapiens. 2005 353 100 0.0 PRO211 polypeptide - Homo sapiens. 2005 353 100 0.0 Protein PRO211 - Homo sapiens. 2005 353 100 0.0 Protein PRO211 - Homo sapiens. 2005 353 100 0.0 Protein PRO211 - Homo sapiens. 2005 353 100 0.0 Protein PRO211 amino acid sequence - Homo s 1997 352 100 0.0 Mature human TANGO 331 protein - Homo sap 1887 329 100 0.0 Unknown (protein for MGC:11256) - Homo sap 1887 329 100 0.0 Human ORFX ORF2475 polypeptide sequence S 1716 298 97 0.0 Human apoptosis related protein - Homo sa 1635 301 85 0.0
sapiens. (353	aa)	nan angiogenesis-associated protein PRO211, SEQ ID NO:57 - Homo [1 seg] [76 bits], Expect = 0.0
		$3/353 \ (100\%)$, Positives = $353/353 \ (100\%)$, at $1,1-353,353$
DNA32292.nc	. 1	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
P_AAB53075	1	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc	61	EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
P_AAB53075	61	EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
DNA32292.nc	121	VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
P_AAB53075	121	VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
DNA32292.nc	181	MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
P_AAB53075	181	MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
DNA32292.nc	241	CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
P_AAB53075	241	CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
DNA32292.nc	301	CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
P_AAB53075	301	CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
Score = 2005	5 (7	an TANGO 331 protein - Homo sapiens. (353 aa) [1 seg] 76 bits), Expect = 0.0 /353 (100%), Positives = 353/353 (100%), at 1,1-353,353
DNA32292.nc	1	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW

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P AAB61231
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
 DNA32292.nc
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
                **********
  P AAB61231
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
                ******************
  P AAB61231
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
  P AAB61231
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 DNA32292.nc
                ****************
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
  P AAB61231
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
 DNA32292.nc
                *******************
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
  P AAB61231
>3 P AAB80212 Human PRO211 protein - Homo sapiens. (353 aa) [1 seg]
 Score = 2005 (776 bits), Expect = 0.0
 Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
 DNA32292.nc
              1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
                ******************
  P AAB80212
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
 DNA32292.nc
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
               ******************
  P AAB80212
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYOGPLCTDC
  P AAB80212
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
               ******************
 P AAB80212
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
               P AAB80212
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
DNA32292.nc
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
               ***************
 P AAB80212
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>4 P_AAB68596 PRO211 - Homo sapiens. (353 aa) [1 seg]
Score = 2005 (776 bits), Expect = 0.0
Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc
               P AAB68596
             1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
```

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DNA32292.nc
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAOEEHLEAWWLOLKSEYPDLFEWFC
               **********
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
  P AAB68596
           121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
               **********
  P AAB68596
           121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
           181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
           181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
  P AAB68596
           241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 DNA32292.nc
               ******************
 P AAB68596
           241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGOCADVDECSLAEKT
 DNA32292.nc
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
               ***********
 P AAB68596
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
>5 P AAY83224 PRO211 Polypeptide - Homo sapiens. (353 aa) [1 seg]
 Score = 2005 (776 bits), Expect = 0.0
 Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
 DNA32292.nc
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
               *****************
 P AAY83224
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
 DNA32292.nc
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
               ******************
 P AAY83224
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAQEEHLEAWWLOLKSEYPDLFEWFC
DNA32292.nc
           121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
               *******************
 P AAY83224
           121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
DNA32292.nc
           181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
               *******************
 P AAY83224
           181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
DNA32292.nc
           241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
               ************
           241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 P AAY83224
DNA32292.nc
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
               P AAY83224
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>6 P_AAB00169 PRO211 polypeptide - Homo sapiens. (353 aa) [1 seg]
Score = 2005 (776 bits), Expect = 0.0
Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
DNA32292.nc
            1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
               P AAB00169
             1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
```

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DNA32292.nc
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAQEEHLEAWWLQLKSEYPDLFEWFC
                ******************
  P AAB00169
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAQEEHLEAWWLOLKSEYPDLFEWFC
 DNA32292.nc 121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
                *******************
  P AAB00169
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYOGPLCTDC
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
                ********************
  P AAB00169
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
                *******************
  P AAB00169
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGOCADVDECSLAEKT
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
 DNA32292.nc
                *****************
  P AAB00169
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>7 P AAY05283 EGF-like homologue PRO211 - Homo sapiens. (353 aa) [1 seg]
 Score = 2005 (776 bits), Expect = 0.0
 Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
 DNA32292.nc
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
                **********************
  P AAY05283
              1 \ \mathsf{MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
 DNA32292.nc
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
 P AAY05283
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
                ***********************
 P AAY05283
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
               *******************
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 P AAY05283
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
               ***********************
 P AAY05283
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 DNA32292.nc
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
               ******************
 P AAY05283
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>8 P AAY13344 protein PRO211 - Homo sapiens. (353 aa) [1 seg]
Score = 2005 (776 bits), Expect = 0.0
Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
DNA32292.nc
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
               *******************
 P AAY13344
            1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc 61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
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P AAY13344
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAOEEHLEAWWLOLKSEYPDLFEWFC
 DNA32292.nc
           121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSROGDGSCRCHMGYOGPLCTDC
                ******************
            121 \ \ VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
  P AAY13344
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
                ***********
  P AAY13344
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
                ***************
  P AAY13344
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 DNA32292.nc
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL .
                ********************
  P AAY13344
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>9 P AAY08064 Human EGF-like homologue protein (PRO217) encoded by DNA32292 cDNA
- Homo sapiens. (353 aa) [1 seq]
 Score = 2005 (776 bits), Expect = 0.0
 Identities = 353/353 (100%), Positives = 353/353 (100%), at 1,1-353,353
 DNA32292.nc
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
               ************************
            1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
  P AAY08064
 DNA32292.nc
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
 P AAY08064
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
               ****************
 P AAY08064
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
               *******************
 P AAY08064
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGOCADVDECSLAEKT
               ******************
 P AAY08064
           241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
DNA32292.nc
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
               ************
 P AAY08064
           301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
>10 P AAY88571 Human PRO211 amino acid sequence - Homo sapiens. (353 aa) [1 seg]
Score = 1997 (773 \text{ bits}), Expect = 0.0
Identities = 352/353 (99%), Positives = 352/353 (99%), at 1,1-353,353
DNA32292.nc
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
               ***************
 P AAY88571
            1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc 61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
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*****************
  P AAY88571
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAQEEHLEAWWLOLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNCHCSGDGSROGDGSCRCHMGYOGPLCTDC
                ******************
  P AAY88571
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
                *****************
  P AAY88571
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGOCADVDECSLAEKT
  P AAY88571
            241 CSAAQFCKMANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
 DNA32292.nc
                ***************
  P AAY88571
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
>11 P AAB61233 Mature human TANGO 331 protein - Homo sapiens. (329 aa) [1 seg]
 Score = 1887 (731 bits), Expect = 0.0
 Identities = 329/329 (100%), Positives = 329/329 (100%), at 25,1-353,329
 DNA32292.nc
             25 AKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAWEEKTLSKYESSEIRLLEILEGLCE
  P AAB61233
              1 AKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAWEEKTLSKYESSEIRLLEILEGLCE
             85 SSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFCVKTLKVCCSPGTYGPDCLACQGGS
 DNA32292.nc
                *****************
  P AAB61233
             61 SSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFCVKTLKVCCSPGTYGPDCLACQGGS
            145 QRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDCMDGYFSSLRNETHSICTACDESCK
 DNA32292.nc
                *****************
  P AAB61233
            121 QRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDCMDGYFSSLRNETHSICTACDESCK
            205 TCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPPCSAAQFCKNANGSYTCEECDSSCV
 DNA32292.nc
                181 TCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPPCSAAQFCKNANGSYTCEECDSSCV
  P AAB61233
 DNA32292.nc
            265 GCTGEGPGNCKECISGYAREHGQCADVDECSLAEKTCVRKNENCYNTPGSYVCVCPDGFE
               *********************
 P AAB61233
            241 GCTGEGPGNCKECISGYAREHGQCADVDECSLAEKTCVRKNENCYNTPGSYVCVCPDGFE
DNA32292.nc
            325 ETEDACVPPAEAEATEGESPTQLPSREDL
               ********
 P AAB61233
           301 ETEDACVPPAEAEATEGESPTQLPSREDL
>12 AAH02894.114 Unknown (protein for MGC:11256) - Homo sapiens (321 aa) [1 seg]
Score = 1770 (686 \text{ bits}), Expect = 0.0
Identities = 321/353 (90%), Positives = 321/353 (90%), Gaps = 32/353 (9%), at
1, 1-353, 321
DNA32292.nc
             1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
               *****************
AAH02894.114
             1 MRLPRRAALGLLPLLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc
            61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAQEEHLEAWWLOLKSEYPDLFEWFC
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AAH02894.114
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAOEEHLEAWWLOLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYOGPLCTDC
                ****************
AAH02894.114
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
                *************************
AAH02894.114
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
 DNA32292.nc
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGOCADVDECSLAEKT
AAH02894.114
            241 CSAAQFCKNANGSYTCE-----DVDECSLAEKT
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
 DNA32292.nc
                ******************
AAH02894.114
            269 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTQLPSREDL
>13 NP 077300.1 hypothetical protein MGC11256 - Homo sapiens (321 aa) [1 seq]
 Score = 1770 (686 \text{ bits}), Expect = 0.0
 Identities = 321/353 (90%), Positives = 321/353 (90%), Gaps = 32/353 (9%), at
1,1-353,321
 DNA32292.nc
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAW
 NP 077300.1
              1 MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
 DNA32292.nc
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
                NP 077300.1
             61 EEKTLSKYESSEIRLLEILEGLCESSDFECNOMLEAOEEHLEAWWLOLKSEYPDLFEWFC
 DNA32292.nc
            121 VKTLKVCCSPGTYGPDCLACOGGSORPCSGNGHCSGDGSROGDGSCRCHMGYOGPLCTDC
               *****************
NP 077300.1
            121 VKTLKVCCSPGTYGPDCLACQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDC
 DNA32292.nc
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
NP 077300.1
            181 MDGYFSSLRNETHSICTACDESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPP
            241 CSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
 DNA32292.nc
               ******
NP 077300.1
            241 CSAAQFCKNANGSYTCE-----DVDECSLAEKT
DNA32292.nc
            301 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
               NP 077300.1
            269 CVRKNENCYNTPGSYVCVCPDGFEETEDACVPPAEAEATEGESPTOLPSREDL
>14 P AAB42711 Human ORFX ORF2475 polypeptide sequence SEQ ID NO:4950 - Homo
(318 aa) [1 seg]
Score = 1716 (665 \text{ bits}), Expect = 0.0
Identities = 298/306 (97%), Positives = 298/306 (97%), at 20,2-325,307
DNA32292.nc
            20 PAPEAAKKPTPCHRCRGLVDKFNOGMVDTAKKNFGGGNTAWEEKTLSKYESSEIRLLEIL
                        ***********
 P AAB42711
             2 PACPPGYLTAPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAWEEKTLSKYESSEIRLLEIL
```

DNA32292.nc	80	EGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFCVKTLKVCCSPGTYGPDCLA
P_AAB42711	62	EGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFCVKTLKVCCSPGTYGPDCLA
DNA32292.nc	140	CQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDCMDGYFSSLRNETHSICTAC ***********************************
P_AAB42711	122	CQGGSQRPCSGNGHCSGDGSRQGDGSCRCHMGYQGPLCTDCMDGYFSSLRNETHSICTAC
DNA32292.nc	200	DESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPPCSAAQFCKNANGSYTCEEC
P_AAB42711	182	DESCKTCSGLTNRDCGECEVGWVLDEGACVDVDECAAEPPPCSAAQFCKNANGSYTCEEC
DNA32292.nc	260	DSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKTCVRKNENCYNTPGSYVCVC
P_AAB42711	242	DSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKTCVRKNENCYNTPGSYVCVC
DNA32292.nc	320	PDGFEE *****
P_AAB42711	302	PDGFEE
Score = 163	5 (63	man apoptosis related protein - Homo sapiens. (353 aa) [1 seg] 34 bits), Expect = 0.0 /354 (85%), Positives = $306/354$ (86%), Gaps = $2/354$ (0%), at
DNA32292.nc	1	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
P AAY91870		*****************
L_M151070	1	**************************************
DNA32292.nc		MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLOLKSEYPDLFEWFC
	61	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW
DNA32292.nc	61 61	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC
DNA32292.nc P_AAY91870	61 61 121	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************
DNA32292.nc P_AAY91870 DNA32292.nc	61 61 121 121	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************
DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870	61 61 121 121 181	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************
DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870 DNA32292.nc	61 61 121 121 181 181	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************
DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870	61 61 121 121 181 181 240	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************
DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870 DNA32292.nc P_AAY91870 DNA32292.nc	61 61 121 121 181 181 240 240	MRLPRRAALGLLPLLLLPPAPEAAKKPTPCHRCRGLVDKFNQGMVDTAKKNFGGGNTAW EEKTLSKYESSEIRLLEILEGLCESSDFECNQMLEAQEEHLEAWWLQLKSEYPDLFEWFC ***********************************